Abstract. Desmoplastic fibroma, alternatively known as aggressive fibromatosis or desmoid tumors, occurs in the form of benign locally aggressive tumors that possess a high rate of recurrence. The forearm bones are rarely involved. The current study presents a case of desmoplastic fibroma in the distal forearm of a 23-year-old man. The tumor was widely resected, and the bone defect was reconstructed using an autologous vascularized fibular graft during the resection procedure. The patient experienced recurrence three times and underwent four resections during the subsequent 3 years following the initial resection. After 10 years of follow-up, the patient's functional recovery remains positive. Despite the implication that surgical resection may be involved in the development of aggressive fibromatosis, surgical wide local excision and functional reconstruction were recommended for the treatment of the present patient.

Introduction

Desmoplastic fibroma, alternatively known as aggressive fibromatosis or desmoid tumors, was initially described by MacFarlane in 1832. The annual incidence of desmoplastic fibroma is 2-4 per 1 million individuals, worldwide (1). Clinically, it presents as a monoclonal fibroblastic proliferation arising in musculoaponeurotic structures (2,3). Magnetic resonance imaging (MRI) is typically used to determine the extent of the tumor and histological analysis is the gold standard for diagnosis. Desmoplastic fibroma can be treated with surgery, radiotherapy or pharmacological agents (1-3). The disease occurs in the form of benign locally aggressive tumors that possess a high rate of recurrence. Desmoplastic fibroma of the bone is rare, and the most common site of occurrence is the jaw, followed by the pelvis and long bones. The forearm bones are rarely involved, however, the development of desmoplastic fibroma following a bone fracture has been previously reported. The majority of studies recommend surgical treatment with a wide free margin resection, followed by reconstruction of the bony defect with a fibular graft (4-7).

The current study presents a case of desmoplastic fibroma in the distal forearm treated by wide resection and reconstructive surgery during the resection procedure. The patient experienced recurrence three times and underwent four resections during the subsequent 3 years following the initial resection. After 10 years of follow-up, the patient's functional recovery remains positive. The present case indicated that repeated recurrences of limb desmoplastic fibroma are not an absolute contraindication for restoration of function, however, frequent follow-up is necessary. Written informed consent was obtained from the patient.

Case report

In October 1999, a 23-year-old man presented at Huashan Hospital, Fudan University (Shanghai, China) with a 2-year history of a progressively swelling tumor in the right forearm following spraining of the right wrist when playing basketball. The patient attended a local hospital (Cixi City, China) for conservative treatment after spraining in 1997. Upon presentation to our hospital, local examination revealed an 8x5-cm solid mass over the palmar and ulnar aspects of the wrist, with an unclear boundary and superficial varicose veins. The overlying skin was higher than normal. The patient demonstrated a full range of motion of the wrist and hand joints. Plain radiograph revealed expansive osteolytic growth involving the right distal radius, and a giant cell tumor of the bone. MRI was performed. Imaging assessment revealed a soft-tissue mass involving the right distal radius, which appeared to be a benign giant cell tumor; however, the possibility of malignancy could not be excluded.

A wide surgical resection of the tumor was performed. The solid mass originated from the distal radius. On gross examination, the encapsulated tumor measured 6x5x3.5 cm. The tumor was excised with the involved radiocarpal joint and ulnar periosteum, proximally up to ~8 cm from the wrist joint. Reconstruction of the bony defect was performed using an autologous vascularized fibular graft. Fibular head and
diaphysis measuring 8 cm were harvested from the ipsilateral leg. The head of the fibula was fitted to match the remaining radiocarpal joint in order to recreate the wrist joint. Distally, the fibular diaphysis was fixed with a radius shaft by a dynamic compression plate in order to reconstruct the forearm and wrist joint. Vessel anastomosis was performed during this procedure, with the dorsal branch of the radial artery anastomosing to the knee arteries and the accompanying vein to the head vein. A biopsy was performed following tumor resection, which resulted in a diagnosis of desmoplastic fibroma. Histologically, the hematoxylin and eosin stained specimen resembled low-grade fibrosarcoma, with banal fibroblasts infiltrating the adjacent tissue in fascicles, without mitosis.

During the 3 years immediately after the tumor resection, the patient experienced recurrence three times, involving the distal ulnar, proximal radius and soft tissue of the dorsal region of the distal forearm, respectively. An additional resection was performed following each recurrence, as well as functional reconstruction following the second and third recurrences. The proximal ulnar and distal radius were fixed by a dynamic compression plate to maintain the length of the forearm, and thumb extension restoration was conducted in order to compensate for the involved extensor hallucis muscle.

Subsequent to 10 years of follow-up, the patient achieved 45-5° flexion-extension of the wrist, and 90-70° pronation-supination and 100-0° flexion-extension of the elbow. The function of the fingers, and the knee and ankle joints was unlimited (Fig. 1).

The patient returned to work, and at that time, MRI revealed no signs of tumor recurrence. In this case, the prognosis was very good and the patient was advised to return every 2 years for further follow-up. However, the patient chose not to return.

Discussion

Desmoplastic fibroma of the bone has been generally defined as a benign tumor that exhibits malignant behavior (8-11). The etiology of desmoplastic fibroma remains to be elucidated. A wide range of factors, including genetic predisposition, estrogen stimulation and traumatic activation, such as fractures and surgery, have been implicated in the development of aggressive fibromatosis (12,13). In the present case, the patient had a history of wrist sprains and had undergone several surgeries, therefore, an association between wrist trauma and tumor growth was suspected, in addition to an association between surgical resections and recurrence.

Due to this locally aggressive behavior, the majority of studies recommend a wide local surgical excision via the normal soft tissue (14-21). The recurrence rate has been reported to be 17% following resection, and may be as high as 55% following curettage (22). In the present case, the patient experienced three further recurrences and underwent four resections.

Wide surgical resection and a high grade of recurrence may result in significant dysfunction of the affected area. Various options have been reported in the relevant literature concerning the functional reconstruction of bone defects in the forearm, including creation of a single forearm bone from the radius and ulna, and centralizing the carpus over the single bone, a free fibular graft, a fibular head graft for the distal radius and a microvascular fibular graft, and allografts (4-7). The technique utilized in the present study to create the articular surface of the radiocarpal joint involved a microvascular fibular head graft. In a total of three surgical procedures, a single bone forearm was created from the radius and ulna, and the carpus was centralized over the single bone. Although a normal forearm could not be achieved, this reconstruction was an acceptable option considering the morbidity of tumor resection for forearm functioning.

The long-term outcome for patients exhibiting extremity desmoid tumors remains to be elucidated, particularly for patients who experience repeated recurrences. In the present case, the patient experienced recurrence three times in the three years immediately after the initial surgical resection, and after 10 years of follow-up, the patient demonstrated positive hand, elbow, knee and ankle function, as well as moderate...
wrist function. The patient was satisfied with the recovery of function and was able to return to work and daily life with no limitations.

In conclusion, the present study indicates that desmoplastic fibroma is a rare tumor that exhibits locally aggressive behavior. The diagnosis is dependent on the correlation of clinico-radio-pathological data. The treatment of choice is a wide local excision and function reconstruction for the upper limbs.

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References